

Konformitätserklärung
Declaration of Conformity

Document-No.: **TD-004**
Revision-No.: **22**
Effective Date: **Last date of signature**
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Wir	We
B. Braun Melsungen AG Carl-Braun-Straße 1 34212 Melsungen Deutschland/Germany	
erklären in eigener Verantwortung, dass das/die Produkt/e	hereby declare in our own responsibility that the product/s
Aqua B. Braun Sterile Water for Irrigation NaCl 0.9% B. Braun Sodium Chloride 0.9% for Irrigation Solución para Irrigación Vitulia NaCl 0,9% 0,9% w/v Sodium Chloride Irrigation Solution Ringer B. Braun Ringer-Lactate B. Braun der Produktgruppe Spüllösungen zur topischen, intra- und postoperativen Verwendung (Artikelnummern siehe Anlage I)	Aqua B. Braun Sterile Water for Irrigation NaCl 0.9% B. Braun Sodium Chloride 0.9% for Irrigation Solución para Irrigación Vitulia NaCl 0,9% 0,9% w/v Sodium Chloride Irrigation Solution Ringer B. Braun Ringer-Lactate B. Braun of the Product Group Solutions for Irrigation for Topical, Intra- and Postoperative use (article numbers see attachment I)
mit den Anforderungen der folgenden Richtlinie übereinstimmt/übereinstimmen	is/are in compliance with the following directive
Richtlinie 93/42/EWG des Rates vom 14. Juni 1993 über Medizinprodukte, geändert durch Richtlinie 2007/47/EG	Council Directive 93/42/EEC of 14 June 1993 concerning Medical Devices, amended by Directive 2007/47/EG
Konformitätsbewertungsverfahren nach Anhang II (ausgenommen Abschnitt 4) der oben genannten Richtlinie	Conformity assessment procedure according to annex II (excluding section 4) of the Directive named above
Klassifizierung gemäß Anhang IX der oben genannten Richtlinie: Klasse IIa	Classification according to annex IX of the Directive named above: Class IIa
Benannte Stelle TÜV SÜD Product Service GmbH Ridlerstraße 65 80339 München Deutschland Kennnummer 0123	Notified body TÜV SÜD Product Service GmbH Ridlerstraße 65 80339 München Germany Identification number 0123
Datum der ersten CE-Kennzeichnung 2003-06	Date of first CE-marking 2003-06
Gültig bis 2024-05-26	Valid until 2024-05-26

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Anlage I / Attachment I

Art.-Nr. / Art. No.	Produktname / Product Name	Klasse / Class
0065729E	Aqua B. Braun	IIa
0066571E	Aqua B. Braun	IIa
0069415E	Aqua B. Braun	IIa
0082423E	Aqua B. Braun	IIa
0082479E	Aqua B. Braun	IIa
3521380	Sterile Water for Irrigation	IIa
3521390	Sterile Water for Irrigation	IIa
3553949	Aqua B. Braun	IIa
3553957	Aqua B. Braun	IIa
3570910	Aqua B. Braun	IIa
387872	Aqua B. Braun	IIa
387873	Aqua B. Braun	IIa
387874	Aqua B. Braun	IIa
387875	Aqua B. Braun	IIa
442464	Aqua B. Braun	IIa
442465	Aqua B. Braun	IIa
442466	Aqua B. Braun	IIa
442467	Aqua B. Braun	IIa
FREU812	Aqua B. Braun	IIa
FREU852	Aqua B. Braun	IIa
FREU912	Aqua B. Braun	IIa
FREU932	Aqua B. Braun	IIa
E30275	Sterile Water for Irrigation	IIa
E30277	Sterile Water for Irrigation	IIa
E70275	Sterile Water for Irrigation	IIa
E70277	Sterile Water for Irrigation	IIa
EB7277	Water for Irrigation	IIa
3637007	Sterile Water for Irrigation	IIa
3637011	Sterile Water for Irrigation	IIa
0066569E	NaCl 0,9% B. Braun	IIa
0066570E	NaCl 0,9% B. Braun	IIa
0069414E	NaCl 0,9% B. Braun	IIa
3521360	Sodium Chloride 0.9% for Irrigation	IIa
3521370	Sodium Chloride 0,9% for Irrigation	IIa
3570100	NaCl 0.9% B. Braun	IIa
3570110	NaCl 0.9% B. Braun	IIa
3570120	NaCl 0.9% B. Braun	IIa
3570130	NaCl 0.9% B. Braun	IIa
3570140	NaCl 0.9% B. Braun	IIa

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3570150	NaCl 0.9% B. Braun	Ila
3570160	NaCl 0.9% B. Braun	Ila
3570170	NaCl 0.9% B. Braun	Ila
3570300	NaCl 0.9% B. Braun	Ila
3570301	Sodium Chloride 0,9% for Irrigation	Ila
3570310	NaCl 0.9% B. Braun	Ila
3570330	NaCl 0,9% B. Braun	Ila
3570340	NaCl 0.9% B. Braun	Ila
3570350	NaCl 0.9% B. Braun	Ila
3570360	NaCl 0,9% B. Braun	Ila
3570370	NaCl 0.9% B. Braun	Ila
3570380	NaCl 0.9% B. Braun	Ila
3570390	NaCl 0,9% B. Braun	Ila
3570410	NaCl 0.9% B. Braun	Ila
3570420	NaCl 0.9% B. Braun	Ila
3570440	NaCl 0.9% B. Braun	Ila
3570450	NaCl 0.9% B. Braun	Ila
3570460	Estericlean NaCl 0,9%	Ila
3570470	NaCl 0.9% B. Braun	Ila
3570480	NaCl 0.9% B. Braun	Ila
3570900	NaCl 0,9% B. Braun	Ila
3637006	Sodium Chloride 0,9% for Irrigation	Ila
3637010	Sodium Chloride 0,9%	Ila
391858	NaCl 0,9% B. Braun	Ila
391859	NaCl 0,9% B. Braun	Ila
391860	NaCl 0,9% B. Braun	Ila
391861	NaCl 0,9% B. Braun	Ila
450268	Solución para Irrigación Vitulia NaCl 0,9%	Ila
450272	Solución para Irrigación Vitulia NaCl 0,9%	Ila
E32275	0,9% w/v Sodium Chloride Irrigation Solution	Ila
E70375	0.9% w/v Sodium Chloride Irrigation Solution	Ila
E70377	0.9% w/v Sodium Chloride Irrigation Solution	Ila
E70477	0,9% w/v Sodium Chloride Irrigation Solution	Ila
497429	NaCl 0,9 %, solution d'irrigation	Ila
FREU850	NaCl 0,9% B. Braun	Ila
FREU910	NaCl 0.9% B. Braun	Ila
FREU930	NaCl 0.9% B. Braun	Ila
FREU950	NaCl 0.9% B. Braun	Ila
FREU970	NaCl 0.9% B. Braun	Ila
3570000	Ringer B. Braun	Ila
3570010	Ringer B. Braun	Ila
3570020	Ringer B. Braun	Ila

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3570030	Ringer B. Braun	IIa
3570040	Ringer B. Braun	IIa
3570050	Ringer B. Braun	IIa
3570060	Ringer B. Braun	IIa
FREU864	Ringer B. Braun	IIa
FREU924	Ringer B. Braun	IIa
FREU944	Ringer B. Braun	IIa
FREU964	Ringer B. Braun	IIa
FREU984	Ringer B. Braun	IIa
3570490	Ringer-Lactate B. Braun	IIa
3570500	Ringer-Lactate B. Braun	IIa
3570510	Ringer-Lactate B. Braun	IIa
3570520	Ringer-Lactate B. Braun	IIa
3570530	Ringer-Lactate B. Braun	IIa
FREU920	Ringer-Lactate B. Braun	IIa

Title: Declaration of Conformity - TD-004 Initiator: Roxana M Gheorghe

This document is signed electronically in compliance with the B. Braun electronic signature policies and procedures by following persons:

UserName: Labeirie, Eva (labeevch)
Title: HC-RA-CH01-Global-Regulatory-Affairs-Manager
Date: Thursday, 23 May 2024, 11:12 W. Europe Daylight Time
Meaning: Document signed as Author

UserName: Ritz, Frank (ritzfrde)
Title: HC-QM DE08 Head QM CoE Pharmaceuticals
Date: Thursday, 23 May 2024, 16:09 W. Europe Daylight Time
Meaning: Approve Document

UserName: Spudy, Bjoern (spudbjde)
Title: HC-RA-DE08P Director GRA Operations CoE Pharma
Date: Friday, 24 May 2024, 07:43 W. Europe Daylight Time
Meaning: Approve Document

UserName: Buenger, Joachim (buenjode)
Title: Director Template & Submission Mgmt
Date: Friday, 24 May 2024, 09:10 W. Europe Daylight Time
Meaning: Approve Document

MANUFACTURER`S DECLARATION

in relation to Regulation (EU) 2023/607 amending Regulations (EU) 2017/745 and (EU) 2017/746 as regards the transitional provisions for certain medical devices and in vitro diagnostic medical devices, in particular with respect to

- the validity of certificates issued under Council Directive 90/385/EEC on Active Implantable Medical Devices (AIMDD) or Council Directive 93/42/EEC on Medical Devices (MDD) (Directive Certificates) *and/or*¹
- the compliance of the devices and us as their manufacturer with the conditions for the continued placing on the market and putting into service

Manufacturer name	B. Braun Melsungen AG
Manufacturer address and contact details	Carl-Braun Straße 1 34212 Melsungen GERMANY
Single Registration Number (SRN) (if available)	DE-MF-000000201

Authorised Representative name (if applicable)	N/A
Authorised Representative address and contact details	N/A
Single Registration Number (SRN) (if available)	N/A

¹ The first condition is not applicable in case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body.

Notified body name (if applicable)	TÜV SÜD Product Service GmbH	☒ See attached schedule
Notified body number (if applicable)	0123	☒ See attached schedule
Directive Certificate number(s) to which this confirmation is made (if applicable)	(1) G1 012974 0607 Rev.02; (2) G1 019717 0032 Rev.00; (3) G1 022239 0080 Rev.03; (4) G2S 012974 0457 Rev.02	☒ See attached schedule
Original expiry date as indicated on the Directive Certificate prior to the extension of the validity (if applicable)	(1) 2024-05-26; (2) 2024-05-26; (3) 2024-05-26; (4) 2024-05-26	☒ See attached schedule
End date of extended validity/transition period	2028-12-31	☒ See attached schedule

We, as the manufacturer declare under our sole responsibility:

- for the above listed **Directive Certificate** (or see attached schedule, if multiple certificates) the conditions for the legal extension of validity as required in Article 120.2 of the MDR are met *and/or*²
- the listed **device(s)** in the attached schedule and we as their manufacturer are in compliance with the conditions listed in Article 120.3c of the MDR for continued placing on the market and putting into service,

namely by fulfilling the following conditions:

- **Directive Certificate(s)** as listed above or in the attached schedule
- Directive Certificate(s) covering the listed device(s) was/were issued after 25 May 2017, was/were valid on 26 May 2021 and have not been withdrawn afterwards.

Choose applicable statements:

☐ Expired *before* 20 March 2023:

² The first condition is not applicable in case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body

- ☐ Before the original date of expiry as indicated on the Directive Certificate(s), we and the notified body have signed written agreement(s) in accordance with Section 4.3, second subparagraph of Annex VII to this Regulation for the conformity assessment(s) in respect of the device(s) covered by the expired certificate(s) or in respect of a device(s) intended to substitute that/those device(s), or
- ☐ A Competent Authority has granted a derogation from the applicable conformity assessment procedure in accordance with Article 59(1) MDR (may be provided upon request), or
- ☐ A Competent Authority has required the manufacturer, in accordance with Article 97(1) MDR, to carry out the applicable conformity assessment procedure (may be provided upon request)

Choose one of the following statements only if a derogation per Article 59(1) or a requirement per Article 97(1) has been granted by a Competent Authority:

- ☐ Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitute(s) and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.
- ☐ We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

- ☒ Expired/expires *after* 20 March 2023:

Choose one applicable statement:

- ☒ Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitute(s) and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.
- ☐ We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

➤ Upclassified devices

In case of devices for which the conformity assessment procedure pursuant to MDD did not require the involvement of a notified body, for which the declaration of conformity was drawn up prior to 26 May 2021 and for which the conformity assessment procedure pursuant to this Regulation requires the involvement of a notified body:

Choose one applicable statement:

- ☐ Formal application(s) to the notified body in accordance with Section 4.3, first subparagraph of Annex VII MDR for conformity assessment has/have been made or will be made/submitted by us to a notified body no later than 26 May 2024 for the device(s) listed in the attached schedule or its/their substitutes and signed written agreement(s) is/will be in place in accordance with Section 4.3, second subparagraph of Annex VII MDR before 26 September 2024.
- ☐ We do not intent to lodge an application for conformity assessment by 26 May 2024, therefore the transition period will end on 26 May 2024.

➤ Quality Management System (QMS)

Choose one applicable statement:

- ☐ A QMS in accordance with Article 10(9) MDR will be put in place by no later than 26 May 2024.
- ☐ A QMS in accordance with Article 10(9) MDR is in place.
- ☒ A notified body has issued the attached certificate for the MDR-compliant QMS.

➤ Device(s) as listed in the attached schedule

- The device(s) continue to comply with the AIMDD or MDD.
- There are no significant changes in the design and intended purpose.
- The device(s) do not present an unacceptable risk to health or safety of patients, users or other persons, or to other aspects of the protection of public health.

Signed for and on behalf of the manufacturer:

	Quality Management	Regulatory Affairs
Full Company Name	B. Braun Melsungen AG	B. Braun Melsungen AG
Location & Date	Melsungen, 2024-07-25	Melsungen, 2024-07-25
Signature	See electronic signature	See electronic signature
Print Name	(1) Thomas Brand; (2) Mareike Arico; (3) Dr. Frank Ritz	(4) Dr. Stefan Seidel; (5) Malte Loh; (6) Dr. Joachim Buenger
Title	(1) Vice President Quality Management for non-active Medical Devices; (2) Head of	(4) Head of Regulatory Affairs CoE Infusion & Pain Therapy; (5) Senior Manager Regulatory



	Quality Management Active Medical Devices/ Head of Regulatory Affairs CoE AIS; (3) Vice President QM Pharma; Hospital Care Division	Affairs; (6) Director Template & Submission Mgmt
Contact Details (at least email)	BBMAG_HC@bbraun.com	BBMAG_HC@bbraun.com
Version of document	Version 4.0	

B. Braun Melsungen AG - Document No.: G10 - Version: 4.0 - Document ID: RE-QM-DIV-000441 - Effective Date: 2024-08-14 - Title: BBMAG_LM_confirmation letter_Regulation EU 2023_607_G10

Schedule of Devices

The above Manufacturer's Declaration is valid for the following devices:

Effective

Identification of the device(s) ³ (e.g., device name, family/group name device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)	Article name (MDR Application)
8717030	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Perfusor® compactplus
8717050	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® compactplus
876100	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	OnlineSuite
8719030	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Spaceplus Perfusor®
8719050	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Spaceplus Infusomat®
8717070	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® compactplus P
4116011F	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Sangofix® Air
4617006	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnifix® Lock

Effective

³ for devices with AIMDD/MDR certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

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932M04SE	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnican fine
931M08SE	NB0123					N/A	Omnican fine
5523606	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drainobag® 600 V
876203	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drug Library Manager Spaceplus
876209	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drug Library Manager Spaceplus
FR29914	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	GLYCINE 1,5 % B. BRAUN
FREU914	NB0123					N/A	GLYCINE 1,5 % B. BRAUN
FREU934						N/A	GLYCINE 1,5 % B. BRAUN
FREU954						N/A	GLYCINE 1,5 % B. BRAUN
FREU974						N/A	GLYCINE 1,5 % B. BRAUN
FREU850	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	NaCl 0,9 % B. BRAUN
FREU910	NB0123					N/A	NaCl 0,9 % B. BRAUN

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FREU930						N/A	NaCl 0,9 % B. BRAUN
FREU950						N/A	NaCl 0,9 % B. BRAUN
FREU970						N/A	NaCl 0,9 % B. BRAUN
3570100	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	NaCl 0,9 % B. BRAUN
3637006						N/A	NaCl 0,9 % B. BRAUN
0069414E						N/A	NaCl 0,9 % B. BRAUN
3521360						N/A	NaCl 0,9 % B. BRAUN
3570120						N/A	NaCl 0,9 % B. BRAUN
3570130						N/A	NaCl 0,9 % B. BRAUN
3570140						N/A	NaCl 0,9 % B. BRAUN
0066570E						N/A	NaCl 0,9 % B. BRAUN
3521370						N/A	NaCl 0,9 % B. BRAUN
3570150						N/A	NaCl 0,9 % B. BRAUN
3570160						N/A	NaCl 0,9 % B. BRAUN
3570170						N/A	NaCl 0,9 % B. BRAUN
0066569E						N/A	NaCl 0,9 % B. BRAUN

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³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

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3570110						N/A	NaCl 0,9 % B. BRAUN
450268	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Vitulia
450272	NB0123					N/A	Vitulia
3570300						N/A	NaCl 0,9 % B. BRAUN
3570301						N/A	NaCl 0,9 % B. BRAUN
3570310						N/A	NaCl 0,9 % B. BRAUN
3570330						N/A	NaCl 0,9 % B. BRAUN
391858						N/A	NaCl 0,9 % B. BRAUN
3570350						N/A	NaCl 0,9 % B. BRAUN
3570360						N/A	NaCl 0,9 % B. BRAUN
3570340						N/A	NaCl 0,9 % B. BRAUN
3637010						N/A	NaCl 0,9 % B. BRAUN
391859						N/A	NaCl 0,9 % B. BRAUN
3570370						N/A	NaCl 0,9 % B. BRAUN
3570380						N/A	NaCl 0,9 % B. BRAUN
3570390						N/A	NaCl 0,9 % B. BRAUN

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³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

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391860						N/A	NaCl 0,9 % B. BRAUN
3570410						N/A	NaCl 0,9 % B. BRAUN
3570420						N/A	NaCl 0,9 % B. BRAUN
497429						N/A	NaCl 0.9% Solution d'irrigation
3570460	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	NaCl 0,9 % B. BRAUN
3570470	NB0123					N/A	NaCl 0,9 % B. BRAUN
3570480						N/A	NaCl 0,9 % B. BRAUN
FREU864	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	RINGER B. BRAUN
FREU920	NB0123					FREU924	RINGER B. BRAUN
FREU944						N/A	RINGER B. BRAUN
FREU964						N/A	RINGER B. BRAUN
FREU984						N/A	RINGER B. BRAUN
3570000	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	RINGER B. BRAUN
3570010	NB0123					N/A	RINGER B. BRAUN
3570020						N/A	RINGER B. BRAUN

Effective

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Schedule of Devices

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3570030						N/A	RINGER B. BRAUN
3570040						N/A	RINGER B. BRAUN
3570050						N/A	RINGER B. BRAUN
3570060						N/A	RINGER B. BRAUN
3570490	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	3570611	RINGER B. BRAUN
3570500	NB0123					3570610	RINGER B. BRAUN
3570510						3570614	RINGER B. BRAUN
3570520						3570612	RINGER B. BRAUN
3570530	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	3570613	RINGER B. BRAUN
FREU812	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Aqua B. Braun
FREU852	NB0123					N/A	Aqua B. Braun
FREU912						N/A	Aqua B. Braun
FREU932						N/A	Aqua B. Braun
387872	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Aqua B. Braun
387873	NB0123					N/A	Aqua B. Braun
387874						N/A	Aqua B. Braun

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442464						N/A	Aqua B. Braun
442465						N/A	Aqua B. Braun
442466						N/A	Aqua B. Braun
3637011						N/A	Sterile Water for Irrigation
3521380	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Aqua B. Braun
3521390	NB0123					N/A	Aqua B. Braun
3553949						N/A	Aqua B. Braun
3553957						N/A	Aqua B. Braun
0065729E						N/A	Aqua B. Braun
0066571E						N/A	Aqua B. Braun
0069415E						N/A	Aqua B. Braun
0082423E						N/A	Aqua B. Braun
0082479E						N/A	Aqua B. Braun
3637007						N/A	Sterile Water for Irrigation
4513800	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Perifix® Catheter Connector
4513801	NB0123					N/A	Perifix® Catheter Connector
4513800N-01						N/A	Perifix® Catheter Connector NRFit
4513801N-01						N/A	Perifix® Catheter Connector NRFit
8713050	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Space

Effective

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	NB0123						
8713070	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Space P
8713030	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Perfusor® Space
8710355	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Enteroport® plus
8700200	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Plus Line SafeSet
8700200-20						N/A	Infusomat® Plus Line SafeSet
8700210						N/A	Infusomat® Plus Line SafeSet
8700310						N/A	Infusomat® Plus Line
8700310-20						N/A	Infusomat® Plus Line
8700310CN						N/A	Infusomat® Plus Line
8250414SP	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Cyto-Set® Infusomat® Space
8250817SP						N/A	Cyto-Set® Infusomat® Space

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8250820SP						N/A	Cyto-Set® Infusomat® Space
8250917SP						N/A	Cyto-Set® Infusomat® Space
8250920SP						N/A	Cyto-Set® Infusomat® Space
835414SP						N/A	Cyto-Set® Infusomat® Space
835817SP						N/A	Cyto-Set® Infusomat® Space
835820SP						N/A	Cyto-Set® Infusomat® Space
835917SP						N/A	Cyto-Set® Infusomat® Space
835920SP						N/A	Cyto-Set® Infusomat® Space
8700420						N/A	Cyto-Set® Infusomat® plus
8700430						N/A	Cyto-Set® Infusomat® plus
8700440						N/A	Cyto-Set® Infusomat® plus
8700450						N/A	Cyto-Set® Infusomat® plus
8700460						N/A	Cyto-Set® Infusomat® plus

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8700470						N/A	Cyto-Set® Infusomat® plus
8700480						N/A	Cyto-Set® Infusomat® plus
8700490						N/A	Cyto-Set® Infusomat® plus
A2581NF	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Cyto-Set® Line
A2582NF	NB0123					N/A	Cyto-Set® Line
A2900N						N/A	Cyto-Set® Mix
A2903N						N/A	Cyto-Set® Mix
A2906N						N/A	Cyto-Set® Mix
A2907N						N/A	Cyto-Set® Mix
A2908N						N/A	Cyto-Set® Mix
4894251	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Stimuplex® A
4894539	NB0123					N/A	Stimuplex® A
4894367						N/A	Stimuplex® A
4894502						N/A	Stimuplex® A
4894375						N/A	Stimuplex® A
4894260						N/A	Stimuplex® A
4894278						N/A	Stimuplex® A
4894278NR						N/A	Stimuplex® A
4894375NR						N/A	Stimuplex® A
4894260NR						N/A	Stimuplex® A
4894367NR						N/A	Stimuplex® A
4894539NR						N/A	Stimuplex® A

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4894502NR						N/A	Stimuplex® A
4894251 NR						N/A	Stimuplex® A
4540002	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Easypump® II LT 60-12
4540002-07	NB0123					N/A	Easypump® II LT 60-12
4540002-20						N/A	Easypump® II LT 60-12
4540003						N/A	Easypump® II LT 500-12.5
4540003-07						N/A	Easypump® II LT 500-12.5
4540003-20						N/A	Easypump® II LT 500-12.5
4540004						N/A	Easypump® II LT 80-16
4540004-07						N/A	Easypump® II LT 80-16
4540004-20						N/A	Easypump® II LT 80-16
4540006						N/A	Easypump® II LT 125-25
4540006-07						N/A	Easypump® II LT 125-25
4540006-20						N/A	Easypump® II LT 125-25
4540008						N/A	Easypump® II LT 270-27
4540008-07						N/A	Easypump® II LT 270-27
4540008-20						N/A	Easypump® II LT 270-27

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4540010						N/A	Easypump® II LT 60-30
4540010-07						N/A	Easypump® II LT 60-30
4540010-20						N/A	Easypump® II LT 60-30
4540012						N/A	Easypump® II LT 120-30
4540012-07						N/A	Easypump® II LT 120-30
4540012-20						N/A	Easypump® II LT 120-30
4540014						N/A	Easypump® II LT 400-40
4540014-07						N/A	Easypump® II LT 400-40
4540014-20						N/A	Easypump® II LT 400-40
4540016						N/A	Easypump® II LT 100-50
4540016-07						N/A	Easypump® II LT 100-50
4540016-20						N/A	Easypump® II LT 100-50
4540018						N/A	Easypump® II LT 270-54
4540018-07						N/A	Easypump® II LT 270-54
4540018-20						N/A	Easypump® II LT 270-54
4540022						N/A	Easypump® II LT 400-80
4540022-07						N/A	Easypump® II LT 400-80

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4540022-20						N/A	Easypump® II LT 400-80
4540026						N/A	Easypump® II LT 270-68
4540026-07						N/A	Easypump® II LT 270-68
4540026-20						N/A	Easypump® II LT 270-68
4540028						N/A	Easypump® II LT 400-100
4540028-07						N/A	Easypump® II LT 400-100
4540028-20						N/A	Easypump® II LT 400-100
4540032						N/A	Easypump® II LT 270-135
4540032-07						N/A	Easypump® II LT 270-135
4540032-20						N/A	Easypump® II LT 270-135
4540040						N/A	Easypump® II ST 100-0,5
4540040-07						N/A	Easypump® II ST 100-0,5
4540040-20						N/A	Easypump® II ST 100-0,5
4540042						N/A	Easypump® II ST 250-0,5
4540042-07						N/A	Easypump® II ST 250-0,5
4540042-20						N/A	Easypump® II ST 250-0,5
4540044						N/A	Easypump® II ST 50-1

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4540044-07						N/A	Easypump® II ST 50-1
4540044-20						N/A	Easypump® II ST 50-1
4540046						N/A	Easypump® II ST 100-1
4540046-07						N/A	Easypump® II ST 100-1
4540046-20						N/A	Easypump® II ST 100-1
4540048						N/A	Easypump® II ST 250-1
4540048-07						N/A	Easypump®II ST 250-1
4540048-20						N/A	Easypump® II ST 250-1
4540050						N/A	Easypump® II ST 250-1,5
4540050-07						N/A	Easypump® II ST 250-1,5
4540050-20						N/A	Easypump® II ST 250-1,5
4540052						N/A	Easypump® II ST 400-2
4540052-07						N/A	Easypump® II ST 400-2
4540052-20						N/A	Easypump® II ST 400-2
4540054						N/A	Easypump® II ST 500-2
4540054-07						N/A	Easypump® II ST 500-2
4540054-20						N/A	Easypump® II ST 500-2

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4540056						N/A	Easypump® II ST 100-2
4540056-07						N/A	Easypump® II ST 100-2
4540056-20						N/A	Easypump® II ST 100-2
4540058						N/A	Easypump® II ST 400-4
4540058-07						N/A	Easypump® II ST 400-4
4540058-20						N/A	Easypump® II ST 400-4
4505000	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	4505000-13	Spinal Introducer
4500059	NB0123					4500059-13	Spinal Introducer
4898650CN	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Contiplex® S 360
4898610CN	NB0123					N/A	Contiplex® S 360
4898615CN						N/A	Contiplex® S 360
4898650-01						N/A	Contiplex® S Ultra 360®
4898610-01						N/A	Contiplex® S Ultra 360®
4898615-01						N/A	Contiplex® S Ultra 360®
4898650-27						N/A	Contiplex® S Ultra 360®
4898610-27						N/A	Contiplex® S Ultra 360®
4898615-27	N/A					Contiplex® S Ultra 360®	

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4515501	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Perifix® Filter
4515501N-01	NB0123					N/A	Perifix® Filter NRFit
4898650NR-27	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Contiplex® S Ultra 360® NRFit®
4898610NR-27	NB0123					N/A	Contiplex® S Ultra 360® NRFit®
4898615NR-27						N/A	Contiplex® S Ultra 360® NRFit®
4898704NR-01						N/A	Contiplex® Tuohy Ultra 360® NRFit®
4898705NR-01						N/A	Contiplex® Tuohy Ultra 360® NRFit®
4898710NR-01						N/A	Contiplex® Tuohy Ultra 360® NRFit®
4898715NR-01						N/A	Contiplex® Tuohy Ultra 360® NRFit®
4898704-01						N/A	Contiplex® Tuohy Ultra 360®
4898705-01						N/A	Contiplex® Tuohy Ultra 360®
4898710-01						N/A	Contiplex® Tuohy Ultra 360®
4898715-01						N/A	Contiplex® Tuohy Ultra 360®

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4898704-27						N/A	Contiplex® Tuohy Ultra 360®
4898705-27						N/A	Contiplex® Tuohy Ultra 360®
4898710-27						N/A	Contiplex® Tuohy Ultra 360®
4898715-27						N/A	Contiplex® Tuohy Ultra 360®
4099117	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Discofix®
4095111	NB0123					N/A	Discofix®
4095120						N/A	Discofix®
4095146						N/A	Discofix®
4095111IN						N/A	Discofix®
409511CN						N/A	Discofix®
409512CN						N/A	Discofix®
16466						N/A	Discofix®
4098102						N/A	Discofix®
409810CN						N/A	Discofix®
4098218						N/A	Discofix®
409821CN						N/A	Discofix®
4098501						N/A	Discofix®
4098234						N/A	Discofix®
4098080						N/A	Discofix®
4055150						N/A	Discofix®
4055145						N/A	Discofix®
4055146						N/A	Discofix®

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4055149						N/A	Discofix®
4055147						N/A	Discofix®
4055148						N/A	Discofix®
4099010						N/A	Discofix®
15809						4095210	Discofix®
9246584	G1 019717 0032 Rev. 00	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	9246605	Nutritub® ENFit® intestinal
9246586	NB0123						
9246576	B. Braun Avitum Italy S.p.A.					9246604	Nutritub® ENFit® intestinal
9246578							
9246519	G1 019717 0032 Rev. 00	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	9246603	Nutritub® Gastral Basic ENFit®
9246518	NB0123					9246602	Nutritub® Gastral Basic ENFit®
9246516	B. Braun Avitum Italy S.p.A.					9246601	Nutritub® Gastral Basic ENFit®
9246550						9246600	Nutritub® Gastral Basic ENFit®
9246515						9246599	Nutritub® Gastral Basic ENFit®
9246592						9246598	Nutritub® Gastral Basic ENFit®
9246514							
9246513							

Effective

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Schedule of Devices

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9246541						9246597	Nutritub® Gastral Basic ENFit®
9246543						9246596	Nutritub® Gastral Basic ENFit®
9246512						9246595	Nutritub® Gastral Basic ENFit®
9246517							
9246525							
9246533						9246594	Nutritub® Gastral Basic ENFit®
9246535							
9246509						9246593	Nutritub® Gastral Basic ENFit®
9246511							
9246508							
8250833SP	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	8250832SP	Infusomat® Space Line
8250835SP	NB0123					8250834SP	Infusomat® Space Line
4238010	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	IN-Stopper
4238011	NB0123					N/A	IN-Stopper
4495101	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Combi-Stopper
4495152	NB0123					N/A	Combi-Stopper
5206634	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Combifix Adapter
5206642	NB0123					N/A	Combifix Adapter

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87229910	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Original Perfusor® Line
4461002	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Pleurofix® No. 1
4461037						N/A	Pleurofix® No. 2
4206096	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Seldinger Introducer Needle
4206100						N/A	Seldinger Introducer Needle
9166432C	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Injekt® 40 Duo
9166432V						N/A	Injekt® 40 Duo
4251127-01	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Introcan Safety® 3
4251127-03						N/A	Introcan Safety® 3
4251127-04						N/A	Introcan Safety® 3
4251127IN						N/A	Introcan Safety® 3
4251127JP						N/A	Introcan Safety® 3
4251128-01						N/A	Introcan Safety® 3
4251128-03						N/A	Introcan Safety® 3

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4251128-04						N/A	Introcan Safety® 3
4251128IN						N/A	Introcan Safety® 3
4251128JP						N/A	Introcan Safety® 3
4251129-01						N/A	Introcan Safety® 3
4251129-03						N/A	Introcan Safety® 3
4251129-04						N/A	Introcan Safety® 3
4251129JP						N/A	Introcan Safety® 3
4251130-01						N/A	Introcan Safety® 3
4251130-03						N/A	Introcan Safety® 3
4251130-04						N/A	Introcan Safety® 3
4251130IN						N/A	Introcan Safety® 3
4251130JP						N/A	Introcan Safety® 3
4251131-01						N/A	Introcan Safety® 3
4251131-03						N/A	Introcan Safety® 3
4251131-04						N/A	Introcan Safety® 3
4251131JP						N/A	Introcan Safety® 3
4251132-01						N/A	Introcan Safety® 3

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4251132-03						N/A	Introcan Safety® 3
4251132-04						N/A	Introcan Safety® 3
4251132IN						N/A	Introcan Safety® 3
4251133-01						N/A	Introcan Safety® 3
4251133-03						N/A	Introcan Safety® 3
4251133-04						N/A	Introcan Safety® 3
4251134-01						N/A	Introcan Safety® 3
4251134-03						N/A	Introcan Safety® 3
4251134-04						N/A	Introcan Safety® 3
4251135-01						N/A	Introcan Safety® 3
4251135-03						N/A	Introcan Safety® 3
4251135-04						N/A	Introcan Safety® 3
4251136-01						N/A	Introcan Safety® 3
4251136-03						N/A	Introcan Safety® 3
4251136-04						N/A	Introcan Safety® 3
4251137-01						N/A	Introcan Safety® 3
4251137-03						N/A	Introcan Safety® 3

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4251137-04						N/A	Introcan Safety® 3
4251144-01						N/A	Introcan Safety® 3
8700036SP	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Space Line
8700435SP	NB0123					N/A	Infusomat® Space Line
8701148SP						N/A	Infusomat® Space Line SafeSet
8270066SP	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	8270066SP-01	Infusomat® Space Line
8270066SP-26	NB0123					N/A	Infusomat® Space Line
8700350-01	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Plus Line
8700350-26	NB0123					N/A	Infusomat® Plus Line
8721748	G1 019717 0032 Rev. 00	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	8721739	Enteroport® ENFit® Set
8721749	NB0123						
8721750	B. Braun Avitum Italy S.p.A.						
8721688							
8721726							
8721734							

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8721735 8721736 8721737 8721742 8721744 8721745 8721746 8721747							
4054032	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Double Spike Adaptor
4094603	NB0123					N/A	Extension Line, Type: Alargadera
4247116						N/A	In-line injection tubing
4053753	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	LS-3 Connector
4097122	NB0123					N/A	LS-2 Connector
4097149						N/A	LS-4 Connector
4097157						N/A	LS-5 Connector
4887441						N/A	Original-Kucher-extension tubing
9500103						N/A	LS-2 Connector

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8250266	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Cyto-Set®
8250366						N/A	ProSet Cyto-Set®
8250370						N/A	ProSet Cyto-Set®
8250455SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250650SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250655SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250818SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250866SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250915SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250966SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250970SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250980SP						N/A	ProSet Cyto-Set® Infusomat® Space

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8250991SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250992SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250993SP						N/A	ProSet Cyto-Set® Infusomat® Space
8250994SP						N/A	ProSet Cyto-Set® Infusomat® Space
8251055SP						N/A	ProSet Cyto-Set® Infusomat® Space
8350866SP						N/A	ProSet Cyto-Set® Infusomat® Space
8350966SP						N/A	ProSet Cyto-Set® Infusomat® Space
8351655SP						N/A	ProSet Cyto-Set® Infusomat® Space
8352055SP						N/A	ProSet Cyto-Set® Infusomat® Space
8352074SP						N/A	ProSet Cyto-Set® Infusomat® Space
8352075SP						N/A	ProSet Cyto-Set® Infusomat® Space

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4182700	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Cyto-Set® Mix
4182701	NB0123					N/A	ProSet Cyto-Set® Mix
4182702						N/A	ProSet Cyto-Set® Mix
4182705						N/A	ProSet Cyto-Set® Mix
4182706						N/A	ProSet Cyto-Set® Mix
4182708						N/A	ProSet Cyto-Set® Mix
4182709						N/A	ProSet Cyto-Set® Line
4182710						N/A	ProSet Cyto-Set® Line
4182711						N/A	ProSet Cyto-Set® Line
4182726						N/A	ProSet Cyto-Set® Mix
4182727						N/A	ProSet Cyto-Set® Mix
4182728						N/A	ProSet Cyto-Set® Line
4182729						N/A	ProSet Cyto-Set® Mix
4182734						N/A	ProSet Cyto-Set® Line
4182817						N/A	ProSet Cyto-Set® Mix
4188090						N/A	ProSet Cyto-Set® Mix

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4188091						N/A	ProSet Cyto-Set® Mix
4188092						N/A	ProSet Cyto-Set® Mix
4188093						N/A	ProSet Cyto-Set® Line
4188925						N/A	ProSet Cyto-Set® Mix
4188926						N/A	ProSet Cyto-Set® Mix
4182704	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Cyto-Set® Pump Adapter
A1673SO	NB0123					N/A	Cyto-Set® Pump Adapter
4037011	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Dosifix®
4037012						N/A	Dosifix®
4037013						N/A	Dosifix®
4037032						N/A	Dosifix®
4037031	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Dosifix®
4033809	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Heidelberger Extension Tubing
4034589						N/A	Heidelberger Extension Tubing
4038703						N/A	Heidelberger Extension Tubing

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4055128						N/A	Heidelberger Extension Tubing
4055136						N/A	Heidelberger Extension Tubing
4097130						N/A	Extension Line, Type: Heidelberger
4097173						N/A	Extension Line, Type: Heidelberger
4097190						N/A	Extension Line, Type: Heidelberger
4097262						N/A	Extension Line, Type: Heidelberger
4097290						N/A	Extension Line, Type: Heidelberger
4097291						N/A	Extension Line, Type: Heidelberger
4097300						N/A	Extension Line, Type: Heidelberger
4097408						N/A	Extension Line, Type: Heidelberger
4055764	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Introcan® Certo
4251300	NB0123					N/A	Introcan® Certo
4251318						N/A	Introcan® Certo

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4251326						N/A	Introcen® Certo
4251334						N/A	Introcen® Certo
4251342						N/A	Introcen® Certo
4251350						N/A	Introcen® Certo
4251369						N/A	Introcen® Certo
4252071B						N/A	Introcen®
4252098B						N/A	Introcen®
4252110B						N/A	Introcen®
4252136B						N/A	Introcen®
4252160B						N/A	Introcen®
4252217B						N/A	Introcen®
4252322B						N/A	Introcen®
4253302						N/A	Introcen®-W Certo
4253310						N/A	Introcen®-W Certo
4253329						N/A	Introcen®-W Certo
4253337						N/A	Introcen®-W Certo
4253345						N/A	Introcen®-W Certo
4253353						N/A	Introcen®-W Certo
4253361						N/A	Introcen®-W Certo
4254074B						N/A	Introcen®-W
4254090B						N/A	Introcen®-W
4254112B						N/A	Introcen®-W

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4254139B						N/A	Introcan®-W
4254171B						N/A	Introcan®-W
4254210B						N/A	Introcan®-W
4254325B						N/A	Introcan®-W
16494CCN	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Discofix® C Safeflow
16495CCN	NB0123					N/A	Discofix® C Safeflow
16501CCN						N/A	Discofix® C Safeflow
16500CCN						N/A	Discofix® C Safeflow
16540CCN						N/A	Discofix® C Safeflow
16520CCN						N/A	Discofix® C Safeflow
4099451	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Intrapur®-Neonat
4093216	NB0123					N/A	Intrapur®
4184637						N/A	Sterifix®
4099354						N/A	Sterifix®
4099303						N/A	Sterifix®
4099257						N/A	Sterifix® Neonat
4099753						4099713	Intrapur®
4099850						4099703	Intrapur® Lipid
4183916						N/A	Intrapur®
4099800						N/A	Intrapur®
4099702						N/A	Intrapur®

Effective

³ for devices with AIMDD/MDR certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)



Schedule of Devices

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4099460						N/A	Intrapur® Neonat Lipid
16500CSF-1	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Discofix® C
16540C						N/A	Discofix® C
16494C						N/A	Discofix® C
16801C						N/A	Discofix® C
16494CSF						N/A	Discofix® C
16800C						N/A	Discofix® C
16504C						N/A	Discofix® C
16501C						N/A	Discofix® C
16760C						N/A	Discofix® C
16495CSF						N/A	Discofix® C
16613C						N/A	Discofix® C
16609C						N/A	Discofix® C
16503C						N/A	Discofix® C
16605C						N/A	Discofix® C
16751C						N/A	Discofix® C
16502C						N/A	Discofix® C
16612C						N/A	Discofix® C
16740C						N/A	Discofix® C
16551CSF						N/A	Discofix® C
16497C						N/A	Discofix® C
16610C						N/A	Discofix® C
16540CSF						N/A	Discofix® C
16720C						N/A	Discofix® C

Effective

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Schedule of Devices

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16520CSF						N/A	Discofix® C
16520C						N/A	Discofix® C
16701C						N/A	Discofix® C
16496C						N/A	Discofix® C
16501CSF-1						N/A	Discofix® C
RU16496C						N/A	Discofix® C
RU16495C						N/A	Discofix® C
CN16496C						N/A	Discofix® C
RU16494C						N/A	Discofix® C
EC16494C						N/A	Discofix® C
CN16494C						N/A	Discofix® C
16611C						N/A	Discofix® C
16608C						N/A	Discofix® C
16600C						N/A	Discofix® C
16501CSF						N/A	Discofix® C
4462556	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Pleuracan®
4462505	NB0123					N/A	Pleuracan® B
4462564	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Pleuracan® Back-Check Valve
5523682	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drainobag® Lock 600

Effective

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16700C	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Discofix® C
16500C						N/A	Discofix® C
16495C						N/A	Discofix® C
16560CSF						N/A	Discofix® C
16901C						N/A	Discofix® C
16615C						N/A	Discofix® C
16560C						N/A	Discofix® C
16494C-01						N/A	Discofix® C
16500CSF						N/A	Discofix® C
16551C						N/A	Discofix® C
16900C						N/A	Discofix® C
BR16496C						N/A	Discofix® C
16614C						N/A	Discofix® C
4052145	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Heidelberger Extension Tubing
4052197						N/A	Heidelberger Extension Tubing
4052197H						N/A	Heidelberger Extension Tubing
4251601-01	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Introcan Safety®
4251601-03						N/A	Introcan Safety®
4251601-04						N/A	Introcan Safety®
4251601JP						N/A	Introcan Safety®
4251607-01						N/A	Introcan Safety®

Effective

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4251607-03						N/A	Introcan Safety®
4251607-04						N/A	Introcan Safety®
4251607JP						N/A	Introcan Safety®
4251614-01						N/A	Introcan Safety®-W
4251614-03						N/A	Introcan Safety®-W
4251614-04						N/A	Introcan Safety®-W
4251614JP						N/A	Introcan Safety®-W
4251620-01						N/A	Introcan Safety®
4251621-01						N/A	Introcan Safety®
4251622-01						N/A	Introcan Safety®
4251623-01						N/A	Introcan Safety®
4251628-01						N/A	Introcan Safety®
4251628-03						N/A	Introcan Safety®
4251628-04						N/A	Introcan Safety®
4251628JP						N/A	Introcan Safety®
4251644-01						N/A	Introcan Safety®
4251644-03						N/A	Introcan Safety®
4251644-04						N/A	Introcan Safety®
4251644JP						N/A	Introcan Safety®
4251652-01						N/A	Introcan Safety®
4251652-03						N/A	Introcan Safety®
4251652-04						N/A	Introcan Safety®
4251652JP						N/A	Introcan Safety®
4251679-01						N/A	Introcan Safety®

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4251679-03						N/A	Introcan Safety®
4251679-04						N/A	Introcan Safety®
4251679JP						N/A	Introcan Safety®
4251687-01						N/A	Introcan Safety®
4251687-03						N/A	Introcan Safety®
4251687-04						N/A	Introcan Safety®
4251687JP						N/A	Introcan Safety®
4251695-01						N/A	Introcan Safety®
4251695-03						N/A	Introcan Safety®
4251695-04						N/A	Introcan Safety®
4251695JP						N/A	Introcan Safety®
4251709-01						N/A	Introcan Safety®
4251709-03						N/A	Introcan Safety®
4251709-04						N/A	Introcan Safety®
4251709JP						N/A	Introcan Safety®
4251717-01						N/A	Introcan Safety®
4251717-03						N/A	Introcan Safety®
4251717-04						N/A	Introcan Safety®
4251890-01						N/A	Introcan Safety®
4251890-03						N/A	Introcan Safety®
4251890-04						N/A	Introcan Safety®
4252500-01						N/A	Introcan Safety®
4252500-03						N/A	Introcan Safety®
4252500-04						N/A	Introcan Safety®
4252519-01						N/A	Introcan Safety®
4252519-03						N/A	Introcan Safety®

Effective

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4252519-04						N/A	Introcan Safety®
4252520-01						N/A	Introcan Safety®
4252527-01						N/A	Introcan Safety®
4252527-03						N/A	Introcan Safety®
4252535-01						N/A	Introcan Safety®
4252535-03						N/A	Introcan Safety®
4252535-04						N/A	Introcan Safety®
4252543-01						N/A	Introcan Safety®
4252551-01						N/A	Introcan Safety®
4252551-03						N/A	Introcan Safety®
4252551-04						N/A	Introcan Safety®
4252560-01						N/A	Introcan Safety®
4252560-03						N/A	Introcan Safety®
4252560-04						N/A	Introcan Safety®
4252578-01						N/A	Introcan Safety®
4252578-03						N/A	Introcan Safety®
4252578-04						N/A	Introcan Safety®
4252586-01						N/A	Introcan Safety®
4252586-04						N/A	Introcan Safety®
4252594-01						N/A	Introcan Safety®
4252594-03						N/A	Introcan Safety®
4252594-04						N/A	Introcan Safety®
4253523-01						N/A	Introcan Safety®-W
4253523-03						N/A	Introcan Safety®-W

Effective

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4253523-04						N/A	Introcan Safety®-W
4253523JP						N/A	Introcan Safety®-W
4253540-01						N/A	Introcan Safety®-W
4253540-03						N/A	Introcan Safety®-W
4253540-04						N/A	Introcan Safety®-W
4253540JP						N/A	Introcan Safety®-W
4253566-01						N/A	Introcan Safety®-W
4253566-03						N/A	Introcan Safety®-W
4253566-04						N/A	Introcan Safety®-W
4253566JP						N/A	Introcan Safety®-W
4253574-01						N/A	Introcan Safety®-W
4253574-03						N/A	Introcan Safety®-W
4253574-04						N/A	Introcan Safety®-W
4253574JP						N/A	Introcan Safety®-W
4253590-01						N/A	Introcan Safety®-W
4253590-03						N/A	Introcan Safety®-W
4253590-04						N/A	Introcan Safety®-W

Effective

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4253604-01						N/A	Introcan Safety®-W
4253604-03						N/A	Introcan Safety®-W
4253604-04						N/A	Introcan Safety®-W
4253604JP						N/A	Introcan Safety®-W
4253612-01						N/A	Introcan Safety®-W
4253612-03						N/A	Introcan Safety®-W
4253612-04						N/A	Introcan Safety®-W
4253639-01						N/A	Introcan Safety®-W
4253639-03						N/A	Introcan Safety®-W
4253639JP						N/A	Introcan Safety®-W
4253639-04						N/A	Introcan Safety®-W
4254503-01						N/A	Introcan Safety®-W
4254503-03						N/A	Introcan Safety®-W
4254503-04						N/A	Introcan Safety®-W
4254511-01						N/A	Introcan Safety®-W
4254511-03						N/A	Introcan Safety®-W
4254511-04						N/A	Introcan Safety®-W

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4254538-01						N/A	Introcan Safety®-W
4254538-03						N/A	Introcan Safety®-W
4254538-04						N/A	Introcan Safety®-W
4254546-01						N/A	Introcan Safety®-W
4254546-03						N/A	Introcan Safety®-W
4254554-01						N/A	Introcan Safety®-W
4254554-03						N/A	Introcan Safety®-W
4254554-04						N/A	Introcan Safety®-W
4254562-01						N/A	Introcan Safety®-W
4254562-03						N/A	Introcan Safety®-W
4254562-04						N/A	Introcan Safety®-W
4254570-01						N/A	Introcan Safety®-W
4254570-03						N/A	Introcan Safety®-W
4254570-04						N/A	Introcan Safety®-W
4254597-01						N/A	Introcan Safety®-W
4254597-03						N/A	Introcan Safety®-W
4254597-04						N/A	Introcan Safety®-W

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4183913	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Intrapur®
4183925						N/A	ProSet Intrapur®
4183926						N/A	ProSet Intrapur®
4183927						N/A	ProSet Intrapur®
4183948						N/A	ProSet Intrapur®
4183949						N/A	ProSet Intrapur®
4184004						N/A	ProSet Intrapur®
4184006						N/A	ProSet Intrapur®
4184007						N/A	ProSet Intrapur®
4184008						N/A	ProSet Intrapur®
4098725						N/A	ProSet Intrapur®
4081002						N/A	ProSet Intrapur®
4099265						N/A	ProSet Sterifix® Neonat
4187822						N/A	ProSet Intrapur®
4184001						N/A	ProSet Intrapur®
4183255						N/A	ProSet Intrapur®
4183245						N/A	ProSet Intrapur®
4183240						N/A	ProSet Intrapur®
4180351						N/A	ProSet Intrapur®
4180350						N/A	ProSet Intrapur®
4188960	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Discofix® C
4188959						N/A	ProSet Discofix® C

Effective

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4188957						N/A	ProSet Discifix® C
4188105						N/A	ProSet Discifix® C
4188071						N/A	ProSet Discifix® C
4187954						N/A	ProSet Discifix® C
4187826						N/A	ProSet Discifix® C
4187202						N/A	ProSet Discifix® C
4187199						N/A	ProSet Discifix® C
4187032						N/A	ProSet Discifix® C
4184963						N/A	ProSet Discifix® C
4184491						N/A	ProSet Discifix® C
4184246						N/A	ProSet Discifix® C
4184030						N/A	ProSet Discifix® C
4184022						N/A	ProSet Discifix® C
4182635						N/A	ProSet Discifix® C
4181234						N/A	ProSet Discifix® C
4180965						N/A	ProSet Discifix® C
4086481						N/A	ProSet Discifix® C

Effective

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4085230						N/A	ProSet Discifix® C
4085213						N/A	ProSet Discifix® C
4187203						N/A	ProSet Discifix® C
4182308						N/A	ProSet Discifix® C
4187527						N/A	ProSet Discifix® C
4180437						N/A	ProSet Discifix® C
4183088						N/A	ProSet Discifix® C
4088698						N/A	ProSet Discifix® C
4084792						N/A	ProSet Discifix® C
4085300SF						N/A	ProSet Discifix® C
4085086						N/A	ProSet Discifix® C
4181027						N/A	ProSet Discifix® C
4184005						N/A	ProSet Discifix® C
4187291						N/A	ProSet Discifix® C
4183312						N/A	ProSet Discifix® C
4185366						N/A	ProSet Discifix® C
4185927						N/A	ProSet Discifix® C

Effective

Effective

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4188188						N/A	ProSet Discifix® C
4086482						N/A	ProSet Discifix® C
4184327						N/A	ProSet Discifix® C
4180439						N/A	ProSet Discifix® C
4180306						N/A	ProSet Discifix® C
4182944						N/A	ProSet Discifix® C
4083255						N/A	ProSet Discifix® C
4187911						N/A	ProSet Discifix® C
4187823						N/A	ProSet Discifix® C
4187878						N/A	ProSet Discifix® C
4085168						N/A	ProSet Discifix® C
4189821						N/A	ProSet Discifix® C
4188958						N/A	ProSet Discifix® C
4187213						N/A	ProSet Discifix® C
4187880						N/A	ProSet Discifix® C
4083254						N/A	ProSet Discifix® C
4189847						N/A	ProSet Discifix® C

Effective

Effective

³ for devices with AIMDD/MDR certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)



Schedule of Devices

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4188198						N/A	ProSet Discifix® C
4183510						N/A	ProSet Discifix® C
4187033						N/A	ProSet Discifix® C
4188072						N/A	ProSet Discifix® C
4183787						N/A	ProSet Discifix® C
4180678						N/A	ProSet Discifix® C
4180679						N/A	ProSet Discifix® C
4187879						N/A	ProSet Discifix® C
4185928						N/A	ProSet Discifix® C
4086879						N/A	ProSet Discifix® C
4188047						N/A	ProSet Discifix® C
4189839						N/A	ProSet Discifix® C
4183852						N/A	ProSet Discifix® C
4185985						N/A	ProSet Discifix® C
4085450SF						N/A	ProSet Discifix® C
4089464						N/A	ProSet Discifix® C
4182737						N/A	ProSet Discifix® C

Effective

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4180300						N/A	ProSet Discifix® C
4183777						N/A	ProSet Discifix® C
4185972						N/A	ProSet Discifix® C
4184521						N/A	ProSet Discifix® C
4182652						N/A	ProSet Discifix® C
4184483						N/A	ProSet Discifix® C
4087930						N/A	ProSet Discifix® C
4184817						N/A	ProSet Discifix® C
4187391						N/A	ProSet Discifix® C
4182720						N/A	ProSet Discifix® C
4185821N						N/A	ProSet Discifix® C
4085434SF						N/A	ProSet Discifix® C
4188225						N/A	ProSet Discifix® C
4186580						N/A	ProSet Discifix® C
4186579						N/A	ProSet Discifix® C
4085500SF						N/A	ProSet Discifix® C
4181778						N/A	ProSet Discifix® C

Effective

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4180459						N/A	ProSet Discifix® C
4188510						N/A	ProSet Discifix® C
4180438						N/A	ProSet Discifix® C
4086945						N/A	ProSet Discifix® C
4187898						N/A	ProSet Discifix® C
4185021						N/A	ProSet Discifix® C
4187529						N/A	ProSet Discifix® C
4088520						N/A	ProSet Discifix® C
4181028						N/A	ProSet Discifix® C
4182638						N/A	ProSet Discifix® C
4088699						N/A	ProSet Discifix® C
4180120						N/A	ProSet Discifix® C
4180677						N/A	ProSet Discifix® C
4182633						N/A	ProSet Discifix® C
4182639						N/A	ProSet Discifix® C
4187838						N/A	ProSet Discifix® C
4084510						N/A	ProSet Discifix® C

Effective

Effective

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4182651						N/A	ProSet Discifix® C
4187834						N/A	ProSet Discifix® C
4180445						N/A	ProSet Discifix® C
4083777						N/A	ProSet Discifix® C
4187308						N/A	ProSet Discifix® C
4184424						N/A	ProSet Discifix® C
4182182						N/A	ProSet Discifix® C
4268091B	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Vasofix® Braunüle®
4268113B	NB0123					N/A	Vasofix® Braunüle®
4268130B						N/A	Vasofix® Braunüle®
4268156B						N/A	Vasofix® Braunüle®
4268172B						N/A	Vasofix® Braunüle®
4268210B						N/A	Vasofix® Braunüle®
4268334B						N/A	Vasofix® Braunüle®
4269071						N/A	Vasofix® Certo
4269098						N/A	Vasofix® Certo
4269110						N/A	Vasofix® Certo
4269136						N/A	Vasofix® Certo

Effective

Effective

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)



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4269152						N/A	Vasofix® Certo
4269179						N/A	Vasofix® Certo
4269217						N/A	Vasofix® Certo
4269225						N/A	Vasofix® Certo
4269330						N/A	Vasofix® Certo
4051807	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Extension Line
4054393	NB0123					N/A	Extension Line
4054394						N/A	Extension Line
4055137						N/A	Extension Line
4055138						N/A	Extension Line
4055139						N/A	Extension Line
4055140						N/A	Extension Line
4090144						N/A	ProSet Extension Line
4090365						N/A	ProSet Spiral Line
4090373						N/A	ProSet Spiral Line
4090381						N/A	ProSet Spiral Line
4090383						N/A	ProSet Spiral Line
4090390						N/A	ProSet Spiral Line
4090438						N/A	ProSet Spiral Line
4091621						N/A	ProSet Extension Line

Effective

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Schedule of Devices

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4091622						N/A	ProSet Extension Line
4091660						N/A	ProSet Extension Line
4268091S-01	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Vasofix® Safety
4268091S-03	NB0123					N/A	Vasofix® Safety
4268113S-01						N/A	Vasofix® Safety
4268113S-03						N/A	Vasofix® Safety
4268130S-01						N/A	Vasofix® Safety
4268130S-03						N/A	Vasofix® Safety
4268156S-01						N/A	Vasofix® Safety
4268156S-03						N/A	Vasofix® Safety
4268172S-01						N/A	Vasofix® Safety
4268172S-03						N/A	Vasofix® Safety
4268210S-01						N/A	Vasofix® Safety
4268210S-03						N/A	Vasofix® Safety
4268334S-01						N/A	Vasofix® Safety
4268334S-03						N/A	Vasofix® Safety
4269071S-01						N/A	Vasofix® Safety
4269071S-03						N/A	Vasofix® Safety
4269071SIN						N/A	Vasofix® Safety
4269071S-20						N/A	Vasofix® Safety
4269098S-01						N/A	Vasofix® Safety
4269098S-03						N/A	Vasofix® Safety
4269098SIN						N/A	Vasofix® Safety
4269098S-20						N/A	Vasofix® Safety

Effective

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4269110S-01						N/A	Vasofix® Safety
4269110S-03						N/A	Vasofix® Safety
4269110SIN						N/A	Vasofix® Safety
4269110S-20						N/A	Vasofix® Safety
4269136S-01						N/A	Vasofix® Safety
4269136S-03						N/A	Vasofix® Safety
4269136SIN						N/A	Vasofix® Safety
4269136S-20						N/A	Vasofix® Safety
4269152S-01						N/A	Vasofix® Safety
4269152S-03						N/A	Vasofix® Safety
4269152S-20						N/A	Vasofix® Safety
4269179S-01						N/A	Vasofix® Safety
4269179S-03						N/A	Vasofix® Safety
4269179SIN						N/A	Vasofix® Safety
4269179S-20						N/A	Vasofix® Safety
4269217S-01						N/A	Vasofix® Safety
4269217S-03						N/A	Vasofix® Safety
4269217S-20						N/A	Vasofix® Safety
4269225S-01						N/A	Vasofix® Safety
4269225S-03						N/A	Vasofix® Safety
4269225S-20						N/A	Vasofix® Safety
4269330S-01						N/A	Vasofix® Safety
4269330S-03						N/A	Vasofix® Safety
4269330S-20						N/A	Vasofix® Safety
4091728	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Spiral Line

Effective

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4091736	NB0123					N/A	ProSet Spiral Line
4091740						N/A	ProSet Spiral Line
4091752						N/A	ProSet Spiral Line
4092539						N/A	ProSet Spiral Line
4092937						N/A	ProSet Spiral Line
4092945						N/A	ProSet Spiral Line
4092953						N/A	ProSet Spiral Line
4092961						N/A	ProSet Spiral Line
4092970						N/A	ProSet Spiral Line
4093054						N/A	ProSet Extension Line
4093115						N/A	ProSet Spiral Line
4093130						N/A	ProSet Spiral Line
4093150						N/A	ProSet Spiral Line
4093170						N/A	ProSet Spiral Line
4093185						N/A	ProSet Spiral Line
4093215						N/A	ProSet Spiral Line
4093230						N/A	ProSet Spiral Line

Effective

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4093250						N/A	ProSet Spiral Line
4093270						N/A	ProSet Spiral Line
4093285						N/A	ProSet Spiral Line
4093402						N/A	ProSet Extension Line
4093437						N/A	ProSet Extension Line
4093585						N/A	ProSet Spiral Line
4093607						N/A	ProSet Spiral Line
4093830						N/A	ProSet Spiral Line
4093850						N/A	ProSet Spiral Line
4093870						N/A	ProSet Spiral Line
4093885						N/A	ProSet Spiral Line
4095251						N/A	ProSet Extension Line
4097531						N/A	ProSet Extension Line
4097572						N/A	Extension Line
4099362						N/A	ProSet Spiral Line
4185841						N/A	ProSet Extension Line
4185842						N/A	ProSet Extension Line

Effective

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4187466						N/A	ProSet Spiral Line
4187467						N/A	ProSet Spiral Line
4187468						N/A	ProSet Spiral Line
4187469						N/A	ProSet Spiral Line
4188080						N/A	ProSet Spiral Line
9500049						N/A	Extension Line
9500057						N/A	Extension Line
9500065						N/A	Extension Line
8700390	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat®plus Line SafeSet
8700391	NB0123					N/A	Infusomat®plus Line SafeSet
8700392	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat®plus Line SafeSet
8700140SP	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Space Line SafeSet
8700141SP	NB0123					N/A	Infusomat® Space Line SafeSet
8700142SP	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Space Line SafeSet

Effective

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4060369L	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Intrafix® Primeline
4060407						N/A	Intrafix® Primeline
4062158						N/A	Intrafix® Primeline
4062158C						N/A	Intrafix® Primeline
4062182						N/A	Intrafix® Primeline
4062955						N/A	Intrafix® Air
4062957E						N/A	Intrafix® Primeline
4062981L						N/A	Intrafix® Primeline
4062982L						N/A	Intrafix® Primeline
4062983L						N/A	Intrafix® Primeline
4063000						N/A	Intrafix® SafeSet
4063001						N/A	Intrafix® SafeSet
4063003						N/A	Intrafix® SafeSet
4063004						N/A	Intrafix® SafeSet
4063004C						N/A	Intrafix® SafeSet
4063004M						N/A	Intrafix® SafeSet
4063005						N/A	Intrafix® SafeSet
4063006						N/A	Intrafix® SafeSet
5524237	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drainobag® Basse Pression

Effective

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5522390	NB0123					N/A	Drainobag® Lock 300
5523753						N/A	Drainobag® 150
5523761						N/A	Drainobag® Lock 150
55237611						N/A	Drainobag® Lock 150
U2000600						5523602	Drainobag® Lock 400
5523605	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drainobag® 600 V
5523648	NB0123					N/A	Drainobag® Lock 600 V
5523649						N/A	Drainobag® Lock 600 V
5524210						N/A	Drainobag® Basse Pression TL
5522322						N/A	Drainobag® 300 V
5522340						N/A	Drainobag® Lock 300 V
55223401						N/A	Drainobag® Lock 300 V
5523702						N/A	Drainobag® 150 V
5523710						N/A	Drainobag® 150 VL
5523729						N/A	Drainobag® Lock 150 V
5523737						N/A	Drainobag® Lock 150 VL

Effective

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55237371						N/A	Drainobag® Lock 150 VL
U2000500						5523601	Drainobag® 400 V
U2000700						5523603	Drainobag® Lock 400 V
5523400	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drainobag® Lock 600 K 10
5523401	NB0123					N/A	Drainobag® Lock 600 K 10
5523427						N/A	Drainobag® Lock 600 K 12
5523428						N/A	Drainobag® Lock 600 K 12
4063144	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Intrafix® SafeSet
4063148	NB0123					N/A	Intrafix® SafeSet
4063287						N/A	Intrafix® Primeline
4088549						N/A	ProSet Intrafix® Primeline
4110000						N/A	Intrafix® SafeSet
4110010						N/A	Intrafix® SafeSet
4180038						N/A	ProSet Intrafix® Primeline
4182001A						N/A	ProSet Intrafix® SafeSet
4182002A						N/A	ProSet Intrafix® SafeSet
4182097						N/A	ProSet Intrafix® SafeSet

Effective

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4182098						N/A	ProSet Intrafix® SafeSet
4182111						N/A	ProSet Intrafix® Primeline
4182179						N/A	ProSet Intrafix® SafeSet
4182409						N/A	ProSet Intrafix® SafeSet
4183450						N/A	ProSet Intrafix® SafeSet
4183455						N/A	ProSet Intrafix® SafeSet
4183665						N/A	ProSet Intrafix® SafeSet
4183791						N/A	ProSet Intrafix® Primeline
4184321						N/A	ProSet Intrafix® SafeSet
4186097						N/A	ProSet Intrafix® SafeSet
4186109						N/A	ProSet Intrafix® SafeSet
4186110						N/A	ProSet Intrafix® SafeSet
4186168						N/A	ProSet Intrafix® Primeline
4186320						N/A	ProSet Intrafix® Primeline
4186711						N/A	ProSet Intrafix® Primeline
4186950						N/A	ProSet Intrafix® Primeline
4186980						N/A	ProSet Intrafix® SafeSet

Effective

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)



Schedule of Devices

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4186981						N/A	ProSet Intrafix® SafeSet
4187005						N/A	ProSet Intrafix® Primeline
4187006						N/A	ProSet Intrafix® SafeSet
4187007						N/A	ProSet Intrafix® Primeline
4187008						N/A	ProSet Intrafix® Primeline
4187009						N/A	ProSet Intrafix® SafeSet
4187010						N/A	ProSet Intrafix® Primeline
4187011						N/A	ProSet Intrafix® SafeSet
4187113						N/A	ProSet Intrafix® SafeSet
4187172						N/A	ProSet Intrafix® Primeline
4187176						N/A	ProSet Intrafix® Primeline
4187334						N/A	ProSet Intrafix® Primeline
4187555						N/A	ProSet Intrafix® Primeline
4187946						N/A	ProSet Intrafix® Primeline
4187989						N/A	ProSet Intrafix® SafeSet
4188020						N/A	ProSet Intrafix® Primeline
4188030						N/A	ProSet Intrafix® SafeSet

Effective

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4188110						N/A	ProSet Intrafix® SafeSet
4188113						N/A	ProSet Intrafix® SafeSet
4188114						N/A	ProSet Intrafix® SafeSet
4188115						N/A	ProSet Intrafix® SafeSet
4188116						N/A	ProSet Intrafix® SafeSet
4188117						N/A	ProSet Intrafix® SafeSet
4187105						N/A	ProSet Intrafix® Primeline
4188120						N/A	ProSet Intrafix® SafeSet
4188136						N/A	ProSet Intrafix® SafeSet
4188137						N/A	ProSet Intrafix® SafeSet
4188140						N/A	ProSet Intrafix® SafeSet
4188155						N/A	ProSet Intrafix® SafeSet
4188159						N/A	ProSet Intrafix® SafeSet
4188170						N/A	ProSet Intrafix® SafeSet
4188530						N/A	ProSet Intrafix® SafeSet
4188531						N/A	ProSet Intrafix® SafeSet
4188540						N/A	ProSet Intrafix® SafeSet

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4188550						N/A	ProSet Intrafix® SafeSet
4189109						N/A	ProSet Intrafix® SafeSet
4189582						N/A	ProSet Intrafix® SafeSet
4188119	G2S 012974 0457 Rev. 02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Intrafix® SafeSet
4062877	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Intrafix® Primeline
4062878						N/A	Intrafix® SafeSet
4110001						N/A	Intrafix® Primeline
4110002						N/A	Intrafix® Primeline
4186914						N/A	ProSet Intrafix®
4060563	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Intrafix® Primeline
4063000A	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	SafeSet
4063001CN						N/A	SafeSet
4063003CN						N/A	SafeSet
4063004CN						N/A	SafeSet
4063004SFCN						N/A	SafeSet
4063005CN						N/A	SafeSet

Effective

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4063006CN						N/A	SafeSet
8700340CN	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Plus Line
8700330CN						N/A	Infusomat® Plus Line
8700240-20						N/A	Infusomat® Plus Line SafeSet
8700280						N/A	Infusomat® Plus Line SafeSet
8700300						N/A	Infusomat® Plus Line SafeSet
8700340						N/A	Infusomat® Plus Line
8700250						N/A	Infusomat® Plus Line SafeSet
8700240						N/A	Infusomat® Plus Line SafeSet
8700220						N/A	Infusomat® Plus Line SafeSet
8700330						N/A	Infusomat® Plus Line
8700320						N/A	Infusomat® Plus Line
4092930	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Original Perfusor® Line
4183945						N/A	ProSet Original Perfusor® Line
4183943						N/A	ProSet Original Perfusor® Line
4183941						N/A	ProSet Original Perfusor® Line

Effective

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4183938						N/A	ProSet Original Perfusor® Line
8723017CN						N/A	Original Perfusor® Line
8722919						N/A	Original Perfusor® Line
8723017						N/A	Original Perfusor® Line
8722919-20						N/A	Original Perfusor® Line
8723017-20						N/A	Original Perfusor® Line
8723018						N/A	Original Perfusor® Line
4183968						N/A	ProSet Original Perfusor® Line
4093000						N/A	ProSet Original Perfusor® Line
4183937						N/A	ProSet Original Perfusor® Line
4183942						N/A	ProSet Original Perfusor® Line
4183947						N/A	ProSet Original Perfusor® Line
4183930						N/A	ProSet Original Perfusor® Line
4183933						N/A	ProSet Original Perfusor® Line
4183935						N/A	ProSet Original Perfusor® Line
4183936						N/A	ProSet Original Perfusor® Line

Effective

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8700350CN	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Plus Line
8700350-20	NB0123					N/A	Infusomat® Plus Line
8700360						N/A	Infusomat® Plus Line
8700132SP	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Space Line
8270074SP	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Space Line
8250908SP	NB0123					N/A	ProSet Infusomat® Space Line
8250902SP						N/A	ProSet Infusomat® Space Line
8250900SP						N/A	ProSet Infusomat® Space Line
8250077SP						N/A	ProSet Infusomat® Space Line
4182586SP						N/A	ProSet Infusomat® Space Line
4181557SP						N/A	ProSet Infusomat® Space Line

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8250958SP						N/A	ProSet Infusomat® Space Line
8700370CN	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Plus Line
8700400	NB0123					N/A	Infusomat® Plus Line
8700370						N/A	Infusomat® Plus Line
9167641WE	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnican® fine
9167650WE	NB0123					N/A	Omnican® fine
9167684WE						N/A	Omnican® fine
9167820WE						N/A	Omnican® fine
929G12S-03						N/A	Omnican® fine
929G12S-41						N/A	Omnican® fine
929G12S-43						N/A	Omnican® fine
931G04S-03						N/A	Omnican® fine
931G04S-41						N/A	Omnican® fine
931G04S-43						N/A	Omnican® fine
931G04SCN						N/A	Omnican® fine
931G04SCN1						N/A	Omnican® fine
931G06S-03						N/A	Omnican® fine
931G06S-41						N/A	Omnican® fine
931G06S-43						N/A	Omnican® fine
931G06S-AP						N/A	Omnican® fine
931G06SCN						N/A	Omnican® fine

Effective

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Schedule of Devices

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931G06SCN1						N/A	Omnican® fine
931G08S-03						N/A	Omnican® fine
931G08S-41						N/A	Omnican® fine
931G08S-43						N/A	Omnican® fine
931G08S-44						N/A	Omnican® fine
932G04S-03						N/A	Omnican® fine
932G04S-41						N/A	Omnican® fine
932G04S-43						N/A	Omnican® fine
932G04S-AP						N/A	Omnican® fine
932G04SCN						N/A	Omnican® fine
932G04SCN1						N/A	Omnican® fine
932G05SCN						N/A	Omnican® fine
932G05SCN1						N/A	Omnican® fine
932G06S-03						N/A	Omnican® fine
932G06S-41						N/A	Omnican® fine
932G06S-43						N/A	Omnican® fine
932G06SCN						N/A	Omnican® fine
932G06SCN1						N/A	Omnican® fine
932P04						N/A	Omnican® fine
932P05						N/A	Omnican® fine
932P06						N/A	Omnican® fine
8700270	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Plus Line SafeSet
8700260-20	NB0123					N/A	Infusomat® Plus Line SafeSet
8700260						N/A	Infusomat® Plus Line SafeSet

Effective

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8722865	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Original Perfusor® Line
8700410	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infusomat® Plus Line
4182190SP	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Infusomat® Space Line
4180639SP						N/A	ProSet Infusomat® Space Line
4180020SP						N/A	ProSet Infusomat® Space Line
8250918SP						N/A	ProSet Infusomat® Space Line
8251001SP						N/A	ProSet Infusomat® Space Line
8251002SP						N/A	ProSet Infusomat® Space Line
4182191SP						N/A	ProSet Infusomat® Space Line
4183900						N/A	ProSet Infusomat® Space Line

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8270058SP						N/A	ProSet Infusomat® Space Line
8252658SP						N/A	ProSet Infusomat® Space Line
8250358SP						N/A	ProSet Infusomat® Space Line
8250903SP						N/A	ProSet Infusomat® Space Line
4182653SP						N/A	ProSet Infusomat® Space Line
4187897						N/A	ProSet Infusomat® Space Line
4184904SP						N/A	ProSet Infusomat® Space Line
4188063SP						N/A	ProSet Infusomat® Space Line
4180635SP						N/A	ProSet Infusomat® Space Line
4188166SP						N/A	ProSet Infusomat® Space Line
4189980SP						N/A	ProSet Infusomat® Space Line

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4186524SP						N/A	ProSet Infusomat® Space Line
4189979SP						N/A	ProSet Infusomat® Space Line
4089340SP						N/A	ProSet Infusomat® Space Line
8250905SP						N/A	ProSet Infusomat® Space Line
4183911						N/A	ProSet Infusomat® Space Line
4185489						N/A	ProSet Infusomat® Space Line
4187769SP						N/A	ProSet Infusomat® Space Line
8251284SP						N/A	ProSet Infusomat® Space Line
4185308SP						N/A	ProSet Infusomat® Space Line
8250904SP						N/A	ProSet Infusomat® Space Line
4186486SP						N/A	ProSet Infusomat® Space Line

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8700095SP						N/A	Infusomat® Space Line
8700110SP						N/A	Infusomat® Space Line
8270350SP						N/A	Infusomat® Space Line
8250710SP						N/A	Infusomat® Space Line
8250731SP						N/A	Infusomat® Space Line
8700131SP						N/A	Infusomat® Space Line
8250719SP						N/A	Infusomat® Space Line
4183878SP						N/A	ProSet Infusomat® Space Line
4180633SP						N/A	ProSet Infusomat® Space Line
8250718SP						N/A	Infusomat® Space Line SafeSet
8700098SP						N/A	Infusomat® Space Line SafeSet
8701149SP						N/A	Infusomat® Space Line SafeSet
8700130SP						N/A	Infusomat® Space Line SafeSet

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8700118SP						N/A	Infusomat® Space Line SafeSet
8250720SP						N/A	Infusomat® Space Line SafeSet
4183918						N/A	ProSet Infusomat® Space Line
4183910						N/A	ProSet Infusomat® Space Line
4187789SP						N/A	ProSet Infusomat® Space Line
4185976SP						N/A	ProSet Infusomat® Space Line
4181558SP						N/A	ProSet Infusomat® Space Line
4089391SP						N/A	ProSet Infusomat® Space Line
8270597SP						N/A	ProSet Infusomat® Space Line
8270358SP						N/A	Infusomat® Space Line SafeSet
4187899						N/A	ProSet Infusomat® Space Line

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4183189SP						N/A	ProSet Infusomat® Space Line
4186940SP						N/A	ProSet Infusomat® Space Line
8700087SP-26						N/A	Infusomat® Space Line
8700087SP-01						N/A	Infusomat® Space Line
8251005SP						N/A	ProSet Infusomat® Space Line
8251004SP						N/A	ProSet Infusomat® Space Line
8251003SP						N/A	ProSet Infusomat® Space Line
4183950SP						N/A	ProSet Infusomat® Space Line
4180631SP						N/A	ProSet Infusomat® Space Line
4183901						N/A	ProSet Infusomat® Space Line
4189981SP						N/A	ProSet Infusomat® Space Line
4187377						N/A	ProSet Infusomat® Space Line

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4182189SP						N/A	ProSet Infusomat® Space Line
8252659SP						N/A	ProSet Infusomat® Space Line
4185687	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Original Perfusor® Line
4085129	NB0123					N/A	ProSet Original Perfusor® Line
8250803						N/A	ProSet Original Perfusor® Line
4183971						N/A	ProSet Original Perfusor® Line
4183970						N/A	ProSet Original Perfusor® Line
8255504N						N/A	Original Perfusor® Line
8745919N						N/A	Original Perfusor® Line
8722940						N/A	Original Perfusor® Line
8723060CN						N/A	Original Perfusor® Line
8255253						N/A	Original Perfusor® Line
8723024						N/A	Original Perfusor® Line
8723023						N/A	Original Perfusor® Line
8723026						N/A	Original Perfusor® Line

Effective

Effective

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)



Schedule of Devices

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8723025						N/A	Original Perfusor® Line
8723021						N/A	Original Perfusor® Line
8723020						N/A	Original Perfusor® Line
8250782						N/A	ProSet Original Perfusor® Line
8250847						N/A	ProSet Original Perfusor® Line
8722941						N/A	Original Perfusor® Line
8722960						N/A	Original Perfusor® Line
8250146						N/A	Original Perfusor® Line
8723060						N/A	Original Perfusor® Line
4185595						N/A	ProSet Original Perfusor® Line
8272565						N/A	Original Perfusor® Line
8255067						N/A	Original Perfusor® Line
8722960-20						N/A	Original Perfusor® Line
8255504NCN						N/A	Original Perfusor® Line
8722862-20						N/A	Original Perfusor® Line
8723060-20						N/A	Original Perfusor® Line
8722862						N/A	Original Perfusor® Line

Effective

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8722935						N/A	Original Perfusor® Line
8255172						N/A	Original Perfusor® Line
8255059						N/A	Original Perfusor® Line
4092933						N/A	ProSet Original Perfusor® Line
4092932						N/A	ProSet Original Perfusor® Line
4092931						N/A	ProSet Original Perfusor® Line
8722935CN						N/A	Original Perfusor® Line
8722870N						N/A	Original Perfusor® Line
8722820						N/A	Original Perfusor® Line
8722935-20						N/A	Original Perfusor® Line
8255490						N/A	Original Perfusor® Line
4183969						N/A	ProSet Original Perfusor® Line
0066088K						N/A	Original Perfusor® Line
0066086H						N/A	Original Perfusor® Line
4180441						N/A	ProSet Original Perfusor® Line
0066087J						N/A	Original Perfusor® Line
0009483H						N/A	Original Perfusor® Line

Effective

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4186850	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	ProSet Infusomat® Space Line
4186842SP						N/A	ProSet Infusomat® Space Line
8700128SP						N/A	Infusomat® Space Line SafeSet
8700127SP						N/A	Infusomat® Space Line
8250437SP						N/A	Infusomat® Space Line
8250438SP						N/A	Infusomat® Space Line SafeSet
8252671SP						N/A	ProSet Infusomat® Space Line
4050192	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Sangofix®
4050192H						N/A	Sangofix®
4050193						N/A	Sangofix®
4052013						N/A	Sangofix®
4052013H						N/A	Sangofix®
4053710						N/A	Sangofix®
4053710H						N/A	Sangofix®
4146492						N/A	Sangofix®
4034228	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Sangofix®

Effective

Effective

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4050151	NB0123					N/A	Sangofix® Air
4051998						N/A	Sangofix®
4051998H						N/A	Sangofix®
4052005						N/A	Sangofix®
4052005H						N/A	Sangofix®
4052218H						N/A	Sangofix®
4080187						N/A	Sangofix® Air
4100514						N/A	Sangofix®
4117301						N/A	Sangofix®
4117549						N/A	Sangofix®
8723001	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Original Perfusor® Line
4094000N	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Infuvalve®
4495209	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Combi-Stopper
4495101R						N/A	Combi-Stopper
4097154N	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Safeflow Extension Set
4097145N						N/A	Safeflow Extension Set
4097154						N/A	Safeflow Extension Set

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409110H	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Safeflow
409100CN	NB0123					N/A	Safeflow
409101H						N/A	Safeflow
409100H						N/A	Safeflow
4097148N	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Safeflow Extension Set
931A04E	G1 012974 0607 Rev. 02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnican® fine
9151117S	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnican® 50
9151125S	NB0123					N/A	Omnican® 50
9151133S						N/A	Omnican® 100
9151141S						N/A	Omnican® 100
9151141SC						N/A	Omnican® 100
9161619S						N/A	Omnican® 20
9161627S						N/A	Omnican® 40
9161627SC						N/A	Omnican® 40
9161635S						N/A	Omnican® 40
9161502S						G1 012974 0607 Rev.02	2024-05-26
9161530S	NB0123	N/A	IBSA FSH/LH				

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16441MS	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Serofine™ needle
16443MS						N/A	Serofine™ needle
16441EMD						N/A	Serofine® needle
16441CA						N/A	B. Braun Pen Needle
P1400060						N/A	Pencylcap™
P1400061						N/A	Pencylcap™
P1400062						N/A	B. Braun Pen Needle
U1244000						N/A	Pencylcap™
U1244100						N/A	Pencylcap®
P1400062CA						N/A	B. Braun Pen Needle
U1244100CA						N/A	B. Braun Pen needle
P1400075						N/A	Pen Needle B. Braun F-Pen DS
16443EMD						N/A	Serofine® needle
5523443	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Drainobag® Lock 600 K 14
5523444						N/A	Drainobag® Lock 600 K 14
5523460						N/A	Drainobag® Lock 600 K 16
5523461						N/A	Drainobag® Lock 600 K 16
5523800						N/A	Drainobag® 150 K 6

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55238001						N/A	Drainobag® 150 K 6
5523850						N/A	Drainobag® 150 K 8
55238501						N/A	Drainobag® 150 K 8
9161333V	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnifix® 40 Duo
9161376C	NB0123					N/A	Omnifix® 100 Duo
9161376V						N/A	Omnifix® 100 Duo
4643011C	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnifix® Luer Duo
4643100V	NB0123					N/A	Omnifix® Luer Duo
4643102C						N/A	Omnifix® Luer Duo
4643102V						N/A	Omnifix® Luer Duo
4643105V						N/A	Omnifix® Luer Duo
4643119C						N/A	Omnifix® Luer Duo
4643119V						N/A	Omnifix® Luer Duo
4643127C						N/A	Omnifix® Luer Duo
4643127V						N/A	Omnifix® Luer Duo
4643135C						N/A	Omnifix® Luer Duo
						N/A	Omnifix® Luer Duo

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4643135V						N/A	Omnifix® Luer Duo
9161465V						N/A	Omnifix®-F Luer Duo
4643161						N/A	Omnifix® Luer Duo
4617022V	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnifix® Luer Lock Solo
4617022V-03	NB0123					N/A	Omnifix® Luer Lock Solo
4617029V						N/A	Omnifix® Luer Lock Solo
4617053V						N/A	Omnifix® Luer Lock Solo
4617053V-03						N/A	Omnifix® Luer Lock Solo
4617100CA						N/A	Omnifix® Luer Lock Solo
4617100V						N/A	Omnifix® Luer Lock Solo
4617100V-03						N/A	Omnifix® Luer Lock Solo
4617207V						N/A	Omnifix® Luer Lock Solo
4617207V-03						N/A	Omnifix® Luer Lock Solo
4617304F						N/A	Omnifix® Luer Lock Solo
4617509F						N/A	Omnifix® Luer Lock Solo
4617509F-03						N/A	Omnifix® Luer Lock Solo

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4617510F-06	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnifix® Luer Lock Solo
4670002S-01	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Sterican® Safety Needle
4670005S-01						N/A	Sterican® Safety Needle
4670008S-01						N/A	Sterican® Safety Needle
4670008SBR						N/A	Sterican® Safety Needle
4670012S-01						N/A	Sterican® Safety Needle
4670016S-01						N/A	Sterican® Safety Needle
4670020S-01						N/A	Sterican® Safety Needle
4670022S-01						N/A	Sterican® Safety Needle
4670025S-01						N/A	Sterican® Safety Needle
4670027S-01						N/A	Sterican® Safety Needle
4670028S-01						N/A	Sterican® Safety Needle
4670030S-01						N/A	Sterican® Safety Needle
4670032S-01						N/A	Sterican® Safety Needle
4670035S-01						N/A	Sterican® Safety Needle

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4670035SBR						N/A	Sterican® Safety Needle
4670040S-01						N/A	Sterican® Safety Needle
4670040SBR						N/A	Sterican® Safety Needle
4670042S-01						N/A	Sterican® Safety Needle
4670045S-01						N/A	Sterican® Safety Needle
4670045SBR						N/A	Sterican® Safety Needle
4670047S-01						N/A	Sterican® Safety Needle
4670050S-01						N/A	Sterican® Safety Needle
4670052S-01						N/A	Sterican® Safety Needle
4670053S-01						N/A	Sterican® Safety Needle
4670055S-01						N/A	Sterican® Safety Needle
4670055SBR						N/A	Sterican® Safety Needle
4650018	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Sterican®
4650034	NB0123					N/A	Sterican®
4657500						N/A	Sterican®
4657519						N/A	Sterican®
4657527						N/A	Sterican®
4657543						N/A	Sterican®

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4657624						N/A	Sterican®
4657640						N/A	Sterican®
4657667						N/A	Sterican®
4657675						N/A	Sterican®
4657683						N/A	Sterican®
4657705						N/A	Sterican®
4657799						N/A	Sterican®
4657853						N/A	Sterican®
4660021						N/A	Sterican®
4665112						N/A	Sterican®
4665120						N/A	Sterican®
4665317						N/A	Sterican®
4665406						N/A	Sterican®
4665457						N/A	Sterican®
4665465						N/A	Sterican®
4665503						N/A	Sterican®
4665511						N/A	Sterican®
4665600						N/A	Sterican®
4665635						N/A	Sterican®
4665643						N/A	Sterican®
4665791						N/A	Sterican®
4667093						N/A	Sterican®
4667123						N/A	Sterican®
9180109						N/A	Sterican®
9180117						N/A	Sterican®
9186158						N/A	Sterican®

Effective

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9186166						N/A	Sterican®
9186174						N/A	Sterican®
9186182						N/A	Sterican®
9166297	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Injekt®-H Luer Duo
4645022C	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Injekt® Luer Duo
4645022UA						N/A	Injekt® Luer Duo
4645022V						N/A	Injekt® Luer Duo
4645057C						N/A	Injekt® Luer Duo
4645057UA						N/A	Injekt® Luer Duo
4645057V						N/A	Injekt® Luer Duo
4645065C						N/A	Injekt® Luer Duo
4645103C						N/A	Injekt® Luer Duo
4645103UA						N/A	Injekt® Luer Duo
4645103V						N/A	Injekt® Luer Duo
4645200C						N/A	Injekt® Luer Duo
4645200UA						N/A	Injekt® Luer Duo
4645200V						N/A	Injekt® Luer Duo
4647220						N/A	Injekt® Luer Duo
9166033V						N/A	Injekt®-F Luer Duo
4670030SBR	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Sterican® Safety Needle

Effective

Effective

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4670053SBR	NB0123					N/A	Sterican® Safety Needle
4898323	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Contiplex® D
4898325						N/A	Contiplex® D
4898305						N/A	Contiplex® D
4898308						N/A	Contiplex® D
4898311						N/A	Contiplex® D
4898335						N/A	Contiplex® D
4898305NR						N/A	Contiplex® D NRFit®
4898335NR						N/A	Contiplex® D NRFit®
4898311NR						N/A	Contiplex® D NRFit®
4898323NR						N/A	Contiplex® D NRFit®
4898325NR						N/A	Contiplex® D NRFit®
4895819NCN						N/A	Contiplex® D
4894235NCN						N/A	Contiplex® D
4894391NCN						N/A	Contiplex® D
4898205	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Contiplex® D
4898211						N/A	Contiplex® D
4898235						N/A	Contiplex® D
4898115	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Contiplex® C

Effective

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

Schedule of Devices

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4898130	NB0123					N/A	Contiplex® C
4898115NR						N/A	Contiplex® C NRFit®
4898130NR						N/A	Contiplex® C NRFit®
4892603-01	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Ultraplex® 360
4892603CN	NB0123					N/A	Ultraplex® 360
4892603NR-01						N/A	Ultraplex® 360 NRFit®
4892605-01						N/A	Ultraplex® 360
4892605CN						N/A	Ultraplex® 360
4892605NR-01						N/A	Ultraplex® 360 NRFit®
4892608-01						N/A	Ultraplex® 360
4892608CN						N/A	Ultraplex® 360
4892608NR-01						N/A	Ultraplex® 360 NRFit®
4892610-01						N/A	Ultraplex® 360
4892610CN						N/A	Ultraplex® 360
4892610NR-01						N/A	Ultraplex® 360 NRFit®
4892615-01						N/A	Ultraplex® 360
4892615CN						N/A	Ultraplex® 360
4892615NR-01						N/A	Ultraplex® 360 NRFit®
4892105	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Stimuplex® D
4892105-23	NB0123					N/A	Stimuplex® D

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4892105CN						N/A	Stimuplex® D
4892105NR						N/A	Stimuplex® D NRFit®
4892108						N/A	Stimuplex® D
4892108-23						N/A	Stimuplex® D
4892108CN						N/A	Stimuplex® D
4892108NR						N/A	Stimuplex® D NRFit®
4892112						N/A	Stimuplex® D
4892112-23						N/A	Stimuplex® D
4892112CN						N/A	Stimuplex® D
4892112NR						N/A	Stimuplex® D NRFit®
4892115						N/A	Stimuplex® D
4892115-23						N/A	Stimuplex® D
4892115NR						N/A	Stimuplex® D NRFit®
4892134						N/A	Stimuplex® D
4892134-23						N/A	Stimuplex® D
4892134NR						N/A	Stimuplex® D NRFit®
4892137						N/A	Stimuplex® D
4892137-23						N/A	Stimuplex® D
4892137NR						N/A	Stimuplex® D NRFit®
4892153						N/A	Stimuplex® D
4892153-23						N/A	Stimuplex® D
4892153NR						N/A	Stimuplex® D NRFit®

Effective

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Schedule of Devices

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Identification of the device(s) ³ (e.g., device name, family/group name device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)	Article name (MDR Application)
4892155						N/A	Stimuplex® D
4892155-23						N/A	Stimuplex® D
4892155NR						N/A	Stimuplex® D NRFit®
4892205						N/A	Stimuplex® D
4892205-23						N/A	Stimuplex® D
4892205NR						N/A	Stimuplex® D NRFit®
4892208						N/A	Stimuplex® D
4892208-23						N/A	Stimuplex® D
4892208NR						N/A	Stimuplex® D NRFit®
4892503-01	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Stimuplex® Ultra 360®
4892503-03	NB0123					N/A	Stimuplex® Ultra 360®
4892503-04						N/A	Stimuplex® Ultra 360®
4892503-20						N/A	Stimuplex® Ultra 360®
4892503CN						N/A	Stimuplex® 360®
4892503NR-01						N/A	Stimuplex® Ultra 360® NRFit®
4892505-01						N/A	Stimuplex® Ultra 360®
4892505-03						N/A	Stimuplex® Ultra 360®
4892505-04						N/A	Stimuplex® Ultra 360®
4892505-20						N/A	Stimuplex® Ultra 360®

Effective

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Schedule of Devices

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Effective

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4892505CN						N/A	Stimuplex® 360®
4892505NR-01						N/A	Stimuplex® Ultra 360® NRFit®
4892508-01						N/A	Stimuplex® Ultra 360®
4892508-03						N/A	Stimuplex® Ultra 360®
4892508-04						N/A	Stimuplex® Ultra 360®
4892508-20						N/A	Stimuplex® Ultra 360®
4892508CN						N/A	Stimuplex® 360®
4892508NR-01						N/A	Stimuplex® Ultra 360® NRFit®
4892510-01						N/A	Stimuplex® Ultra 360®
4892510-03						N/A	Stimuplex® Ultra 360®
4892510-04						N/A	Stimuplex® Ultra 360®
4892510-20						N/A	Stimuplex® Ultra 360®
4892510CN						N/A	Stimuplex® 360®
4892510NR-01						N/A	Stimuplex® Ultra 360® NRFit®
4892515-01						N/A	Stimuplex® Ultra 360®
4892515-03						N/A	Stimuplex® Ultra 360®
4892515-04						N/A	Stimuplex® Ultra 360®
4892515-20						N/A	Stimuplex® Ultra 360®

Effective

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

Schedule of Devices

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Identification of the device(s) ³ (e.g., device name, family/group name device model or catalogue number)	Directive Certificate number(s) to which this confirmation is made (if applicable)	Original expiry date as indicated on the Directive Certificate (s) prior to the extension of the validity (if applicable)	Notified Body name and number that issued the Directive Certificate (if applicable)	Notified Body name and number where the MDR application was lodged/contract signed (if applicable)	End date of extended validity / transition period	Substitute Device(s) (if applicable)	Article name (MDR Application)
4892515CN						N/A	Stimuplex® 360®
4892515NR-01						N/A	Stimuplex® Ultra 360® NRFit®
4617003	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Omnifix® Lock
4617014	NB0123					N/A	Omnifix® Lock
4617021						N/A	Omnifix® Lock
4617508F-01						N/A	Omnifix® Lock
8728615						NB0123	N/A
8728615C	Original Perfusor® Syringe 20 ml						
8728623	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Original Perfusor® Syringe 20 ml
8728623C	NB0123					N/A	Original Perfusor® Syringe 20 ml
8728810F-04						N/A	Original Perfusor® Syringe 50 ml
8728810F						8728810F-06	Original Perfusor® Syringe 50 ml
8728810F-20						N/A	Original Perfusor® Syringe 50 ml
8728844F-04						G1 012974 0607 Rev.02	2024-05-26

Effective

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Schedule of Devices

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8728844F	NB0123					8728844F-06	Original Perfusor® Syringe 50 ml
8728844F-20						N/A	Original Perfusor® Syringe 50 ml
8728852F-04	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Original Perfusor® Syringe 50 ml
8728852F-06	NB0123					N/A	Original Perfusor® Syringe 50 ml
8728852F-20						N/A	Original Perfusor® Syringe 50 ml
8728861F-04	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Original Perfusor® Syringe 50 ml
8728861F-06	NB0123					N/A	Original Perfusor® Syringe 50 ml
8728861F-20						N/A	Original Perfusor® Syringe 50 ml
8728845F-01	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Original Perfusor® Syringe 50 ml
	NB0123						
4450100	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Cystofix®
4450120	NB0123					N/A	Cystofix®
4450130						N/A	Cystofix®

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Schedule of Devices

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4450150						N/A	Cystofix®
4450160						N/A	Cystofix®
4450170						N/A	Cystofix®
4450180						N/A	Cystofix®
4450200	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Cystofix®
4450220	NB0123					N/A	Cystofix®
4450410	G1 022239 0080 Rev. 03	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Cystofix SG
4450412	NB0123					N/A	Cystofix SG
4450414	B.BRAUN MEDICAL SAS					N/A	Cystofix SG
4450416						N/A	Cystofix SG
4450010	G1 022239 0080 Rev. 03	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Cystofix
4450012	NB0123					N/A	Cystofix
4450014	B.BRAUN MEDICAL SAS		TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)		N/A	Cystofix
4450016						N/A	Cystofix
4450512						N/A	Cystofix
4450514						N/A	Cystofix
4450516						N/A	Cystofix
4450712						N/A	Cystofix
4450714						N/A	Cystofix
4450716						N/A	Cystofix

Effective

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4450718						N/A	Cystofix
4450720						N/A	Cystofix
6031510	G1 012974 0607 Rev.02	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	N/A	Vasco® OP Powdered
6031525	NB0123					N/A	Vasco® OP Powdered
6031532						N/A	Vasco® OP Powdered
6031546						N/A	Vasco® OP Powdered
6031553						N/A	Vasco® OP Powdered
6031564						N/A	Vasco® OP Powdered
6080990						N/A	Vasco® OP Sensitive
6081002						N/A	Vasco® OP Sensitive
6081010						N/A	Vasco® OP Sensitive
6081029						N/A	Vasco® OP Sensitive
6081037						N/A	Vasco® OP Sensitive
6081045						N/A	Vasco® OP Sensitive
6081053						N/A	Vasco® OP Sensitive
6081060						N/A	Vasco® OP Sensitive
6081199						N/A	Vasco® OP Underglove

Effective

Effective

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6081200						N/A	Vasco® OP Underglove
6081218						N/A	Vasco® OP Underglove
6081226						N/A	Vasco® OP Underglove
6081234						N/A	Vasco® OP Underglove
6081242						N/A	Vasco® OP Underglove
6081259						N/A	Vasco® OP Underglove
6081267						N/A	Vasco® OP Underglove
6081308						N/A	Vasco® OP eco
6081316						N/A	Vasco® OP eco
6081324						N/A	Vasco® OP eco
6081332						N/A	Vasco® OP eco
6081340						N/A	Vasco® OP eco
6081359						N/A	Vasco® OP eco
6081367						N/A	Vasco® OP eco
6081375						N/A	Vasco® OP eco
6081409						N/A	Vasco® OP Grip
6081417						N/A	Vasco® OP Grip
6081425						N/A	Vasco® OP Grip
6081433						N/A	Vasco® OP Grip
6081441						N/A	Vasco® OP Grip
6081450						N/A	Vasco® OP Grip
6081468						N/A	Vasco® OP Grip

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6081476						N/A	Vasco® OP Grip
9208291						N/A	Vasco® OP Free
9208305						N/A	Vasco® OP Free
9208313						N/A	Vasco® OP Free
9208321						N/A	Vasco® OP Free
9208330						N/A	Vasco® OP Free
9208348						N/A	Vasco® OP Free
9208356						N/A	Vasco® OP Free
9208364						N/A	Vasco® OP Free
U2170701	G1 012974 0607 Rev.02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	5524913	Drainobag® Connection Tube Bayonet
4550404	G2S 012974 0457 Rev. 02 NB0123	2024-05-26	TÜV SÜD Product Service GmbH (NB0123)	TÜV SÜD Product Service GmbH (NB0123)	2028-12-31	415040	Filter Needle
4551001						418021	Filter Hub
4550200						415020	Filter Straw
4550250						415021	Filter Straw
4550200N						339171	Sterifix® Filter Straw 4"
4550250N						339170	Sterifix® Filter Straw 1.75"
4550404N						339169	Sterifix® Filter Needle 1.5"

Effective

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Effective

Document History

Version	Description of Change
1.0	Initial version
2.0	Revision numbers of MDD certificates were added
3.0	Removed BUDI number, article name moved to the end of the table, article number changed for identification of device and substitute device, added article 497429 and 931A04E
4.0	Changed contact details e-mail address

Effective

³ for devices with AIMDD/MDD certificate(s) the identification should be as in the certificate, and only if the certificate has a generic scope it should be as defined above)

This document is signed electronically in compliance with the B. Braun electronic signature policies and procedures by following persons:

UserName: Seidel, Stefan (seidstde)
Title: Vice President Regulatory Affairs CoE Infusion & Pain Therapy
Date: Wednesday, 31 July 2024, 17:11 W. Europe Daylight Time
Meaning: Document signed as Author

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UserName: Seidel, Stefan (seidstde)
Title: Vice President Regulatory Affairs CoE Infusion & Pain Therapy
Date: Wednesday, 31 July 2024, 17:20 W. Europe Daylight Time
Meaning: Approve Document

=====

UserName: Buenger, Joachim (buenjode)
Title: Director Template & Submission Mgmt
Date: Thursday, 01 August 2024, 06:59 W. Europe Daylight Time
Meaning: Approve Document

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UserName: Ritz, Frank (ritzfrde)
Title: HC-QM DE08 Head QM CoE Pharmaceuticals
Date: Thursday, 01 August 2024, 08:50 W. Europe Daylight Time
Meaning: Approve Document

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UserName: Brand, Thomas (brantode)
Title: HC-QM-DE08 Vice President QM for non-active Medical Devices
Date: Thursday, 01 August 2024, 10:00 W. Europe Daylight Time
Meaning: Approve Document

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Title: BBMAG_LM_confirmation letter_Regulation EU 2023_607_G10 Initiator: Anja Mai

This document is signed electronically in compliance with the B. Braun electronic signature policies and procedures by following persons:

UserName: Arico, Mareike (sommrde)
Title: HC-QM - Head of QM active MD/ Head of Regulatory Affairs CoE AIS
Date: Friday, 02 August 2024, 11:21 W. Europe Daylight Time
Meaning: Approve Document

=====

UserName: Loh, Malte (lohmatde)
Title: HC-RA-DE08 Senior Manager Regulatory Affairs
Date: Monday, 12 August 2024, 10:45 W. Europe Daylight Time
Meaning: Approve Document

=====

UserName: Meyer, Frank (meyefrde)
Title: HC-QM-DE08 Vice President QM Applications Hospital Care
Date: Wednesday, 14 August 2024, 13:06 W. Europe Daylight Time
Meaning: Final Release of the Document

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